

GENETICS AND GENOMICS DIAGNOSTIC LABORATORY

For local courier service and/or inquiries, please contact 513-636-4474 • Fax: 513-636-4373 www.cincinnatichildrens.org/diagnosticlabs • Email: labgeneticcounselors@cchmc.org

Shipping Address:

3333 Burnet Avenue, Room R1042 Cincinnati, OH 45229-3039

Deliveries accepted Monday-Saturday

PEDIATRIC/ADULT GENETIC TEST REQUISITION

All Information Must Be Completed Before Sample Can Be Processed

PATIENT INFORMATION	SPECIMEN INFORMATION		
Patient Name:,,,,,,,	SPECIMEN TYPE: ☐ Peripheral blood ☐ Skin biopsy (site):		
Home Phone: MR# Date of Birth / / Sex Assigned at Birth: Male Female Uncertain/Other:	Specimen Date: / / Time: Specimen Amount: DRAWN BY:		
INDICATIONS/DIAG	SNOSIS/ICD-10 CODE		
□ ADD/ADHD □ Acute myelogenous leukemia (AML) □ Hydrocephalus □ Amenorrhea: 1' or 2' □ Hyper/Hypopigmentation □ Aplastic Anemia □ Autism Spectrum Disorder □ Immune deficiency □ Broad thumbs and/or halluces □ Intellectual disability □ Congenital heart anomaly □ Developmental Delay □ Developmental Delay □ Dysmorphic features □ MRI, abnormal □ Encephalopathy □ Eye anomaly □ Microcephaly □ Erythematous "butterfly" lesion on face □ Myelodysplastic syndrome (MDS)	□ PDD-NOS □ Pancytopenia □ Seizures, convulsions □ Short stature □ Other: □ Newborn Indications: □ Abnormal NIPT/prenatal screen □ Suspected trisomy 21 □ Suspected Turner's syndrome □ Ambiguous genitalia □ Other: □ Consanguinity (describe relationship): □ Known Chromosome Abnormality:		
BILLING INFORMATION (Choose ONE payment method)	PROVIDER INFORMATION		
REFERRING INSTITUTION Institution: Address: City/State/Zip: Accounts Payable Contact Name: Phone: Fax: Email:	Provider Name (print):		
COMMERCIAL INSURANCE* Insurance can only be billed if requested at the time of service. Policy Holder Name: Gender: Date of Birth // Authorization Number: Insurance ID Number:	Referring Physician Signature (REQUIRED) Contact information for results/questions (if different than ordering provider): Name and Title: Phone: () Fax: () Email:		
Insurance Name:	ETHNIC/RACIAL BACKGROUND (Choose All)		
Insurance Address: City/State/Zip: Insurance Phone Number: * PLEASE NOTE: * We will not bill Medicaid, Medicaid HMO, or Medicare except for the following: CCHMC Patients, CCHMC Providers, or Designated Regional Counties. * If you have questions, please call 1-866-450-4198 for complete details.	□ European American (White) □ African-American (Black) □ Native American or Alaskan □ Asian-American □ Pacific Islander □ Ashkenazi Jewish ancestry □ Latino-Hispanic □ (specify country/region of origin) □ Other □ (specify country/region of origin)		

☐ Patient signed completed ABN

Medical Necessity Regulations: At the government's request, the Molecular Genetics Laboratories would like to remind all physicians that when ordering tests that will be paid under federal health care programs, including Medicare and Medicaid programs, that these programs will pay only for those tests the relevant program deems to be (1) included as covered services, (2) reasonable, (3) medically necessary for the treatment and diagnosis of the patient, and (4) not for screening purposes.



REQUIRED: Patient Name: Date of Birth:
--

TEST(S) REC	QUESTED
Cytogenetic Testing	Fanconi Anemia Testing
Chromosome Analysis	☐ Fanconi Anemia (FA) Chromosome Breakage Study
☐ Routine chromosome analysis*	☐ Fanconi Anemia Gene Sequencing Panel
☐ Chromosome mosaicism study*	(BRCA1, BRCA2, BRIP1, ERCC4, FANCA, FANCB, FANCC, FANCD2, FANCE,
☐ Reflex to SNP Microarray if chromosome results are normal [†]	FANCF, FANCG, FANCI, FANCL, MAD2L2, PALB2, RAD51, RAD51C, RFWD3, SLX4, UBE2T, XRCC2)
☐ High resolution chromosome analysis*	If both FA Breakage Study and FA Gene Seq Panel are ordered, testing will
☐ Chromosome mosaicism study*	be run sequentially (breakage study then molecular sequencing if breakage
☐ Reflex to SNP Microarray if chromosome results are normal [†]	study is <u>positive</u> ; if breakage study is negative, molecular sequencing wing the performed by unless concurrent testing is selected here:
*For chromosome analysis: reflex STAT prelim results on infants <1 month. Reflex	☐ Concurrent FA testing is requested
to mosaicism study when sex chromosome/mosaic aneuploidy abnormality	Single Gene Sequencing
suspected by laboratory based on indications provided.	☐ FANCA full gene sequencing ☐ FANCG full gene sequencing
[†] Additional charge for reflex testing. If SNP Microarray is denied by insurance, Chromosome Analysis will be performed as the first test in the algorithm.	□ FANCE full gene sequencing
Microarray	Molecular Genetic Testing
☐ SNP Microarray - Constitutional	☐ ABCD1 gene sequencing (X-Linked Adrenoleukodystrophy)
☐ Episignature Complete Analysis	□ Reflex to <i>ABCD1</i> deletion/duplication by MLPA
☐ Episignature Targeted Analysis: Specify episignature [†] :	☐ ABCD1 deletion/duplication by MLPA
†Please see Episignature Analysis test information sheet for available conditions	☐ Cleft and Craniofacial Gene Panel (288 genes)
	ABCC9, ACSS2, ACTB, ACTG1, ADAMTSL4, AHDC1, ALPL, ALX1, ALX3, ALX4
Optical Genome Mapping	AMELX, AMER1, AMMECR1, AMOTL1, ANKH, ANKRD11, ARHGAP29, ARSE ASPH, ASXL1, ASXL3, B3GAT3, B3GLCT, BCOR, BMP2, BMP4, BMPR1B
☐ Optical Genome Mapping (Genome-wide)	BPNT2, BRAF, BRD4, C2CD3, CBFB, CCNQ, CD96, CDC45, CDH1, CDKN1C
☐ Optical Genome Mapping [†] - Targeted Analysis : Known SV, gene and/or	CDON, CENPF, CEP164, CHD5, CHD7, CILK1, CNOT1, COG1, COL11A1
specific region:	COL11A2, COL2A1, COL9A1, COL9A2, COL9A3, COLEC10, COLEC11 CPLANE1, CREBBP, CTNND1, CTSK, CYP26B1, DDX59, DHCR7, DHODH
†Please contact GGDL to confirm OGM's coverage for the target region before ordering	DISP1, DLL1, DLX4, DPF2, DPH1, DVL1, DVL3, EDN1, EDNRA, EFNA4, EFNB1 EFTUD2, EHMT1, EIF4A3, EP300, ERF, ESCO2, ESRP2, EVC, EVC2, EYA1 FAM20C, FBN1, FGD1, FGF10, FGF8, FGF9, FGFR1, FGFR2, FGFR3, FLNA
FISH (Fluorescent In Situ Hybridization)	FLNB, FOXE1, FOXI3, FRAS1, FREM1, FST, FTO, FZD2, GAS1, GDF11, GJA1
☐ 22q11.2 del (VCFS) (metaphase FISH)	GLI2, GLI3, GNAI3, GNAS, GNPTAB, GPC3, GPC4, GRHL3, GSC, GTF2E2
☐ SRY (Xp11.1q11.1/Yp11.2) (metaphase FISH)	GZF1, HDAC8, HIST1H1E, HNRNPK, HUWE1, HYAL2, HYLS1, IDS, IDUA IFT122, IFT140, IFT43, IGF1R, IGF2, IHH, IL11RA, INPPL1, IRF6, IRX5, ISM1
☐ X/Y centromeres (Xp11.1q1.1/Yp11.1q11.1) (interphase FISH)	JAG1, KAT6A, KAT6B, KDM1A, KDM6A, KIAA0586, KIF7, KMT2D, KRAS
☐ Other FISH (please call lab):	LOXL3, LRP2, LTBP1, MAFB, MAP3K7, MASP1, MED13L, MED25, MEGF8 MEIS2, MID1, MKS1, MN1, MSX1, MSX2, MTX2, MYCN, MYMK, MYT1, NBAS
	NECTIN1, NEDD4L, NIPBL, OFD1, P4HB, PAX1, PAX3, PAX7, PDE4D, PGM1
Other Testing	PHEX, PHF21A, PHF8, PIEZO2, PIGN, PJA1, PLCB4, PLCH1, PLEKHA5 PLEKHA7, PLOD3, POLR1A, POLR1B, POLR1C, POLR1D, POR, PORCN
☐ Special study:	PPP1R12A, PRRX1, PSAT1, PTCH1, PTDSS1, PTPN11, RAB23, RAD21, RAX
☐ Cell Culture, storage & freezing	RBM10, RECQL4, RIPK4, ROR2, RPGRIP1L, RPL5, RSPRY1, RUNX2, RYK SATB2, SCARF2, SCLT1, SCN4A, SEC24D, SEMA3E, SF3B2, SF3B4, SHH
□ Other:	SHOC2, SHROOM3, SIN3A, SIX1, SIX2, SIX3, SIX5, SKI, SLC25A24, SMAD2
	SMAD3, SMAD4, SMAD6, SMARCA4, SMARCB1, SMC1A, SMC3, SMG9, SMC SMS, SMURF1, SNRPB, SON, SOST, SOX11, SOX6, SOX9, SPECC1L, SPRY1
Cytogenetic and Molecular Genetic Testing	SPRY4, STAG2, STIL, SUFU, SUMO1, TBC1D32, TBX1, TBX22, TCF12, TCOF1 TFAP2A, TFAP2B, TGDS, TGFB1, TGFB2, TGFB3, TGFBR1, TGFBR2, TGIF1
Neurodevelopmental Reflex Genetic Test**	TLK2, TMCO1, TOPORS, TP63, TRAF7, TRRAP, TWIST1, TWIST2, TXNL4A
Tests will be run sequentially based on your selection below:	UBE3B, USP9X, VAX1, VCAN, WASHC5, WDR19, WDR35, WNT5A, YAP1 YWHAE, ZEB2, ZIC1, ZIC2, ZNF462, ZSWIM6
☐ Patient is macrocephalic: $\underline{SNP\ Microarray} \rightarrow \underline{Fragile\ X} \rightarrow \underline{PTEN}$	☐ REFLEX to Whole Exome Sequencing th (See additional details below
☐ Male patient with normal or small head circumference:	
SNP Microarray → Fragile X ☐ Female patient with normal or small head circumference:	☐ DNA Extraction and Storage
·	☐ Fragile X DNA testing
SNP Microarray → Fragile X → MECP2	☐ MECP2 sequence analysis (Rett syndrome)
**If SNP Microarray is denied by insurance, Chromosome Analysis will be performed as the first test in the algorithm. See page 3 for additional information.	☐ MECP2 deletion/duplication analysis by MLPA
Chuamanana Buahana Biandara Tastina	\square Prader-Willi/Angelman - by methylation-sensitive MLPA
Chromosome Breakage Disorders Testing	☐ PTEN Autism Spectrum Disorder sequencing
☐ Bloom Syndrome - Sister Chromatid Exchange (SCE) analysis	☐ Rubinstein-Taybi and Related Syndromes Gene Panel
☐ Chromosome Breakage Disorders Gene Sequencing Panel	(CREBBP, EP300, HNRNPH1, HNRNPH2, SIN3A, SIN3B, SRCAP with CREBBP and EP300 deletion/duplication analysis by MLPA)
(ATM, BLM, BRCA1, BRCA2, BRIP1, ERCC4, FANCA, FANCB, FANCC, FANCD2, FANCE, FANCF, FANCG, FANCI, FANCL, LIG4, MAD2L2, MYSM1, NBN, NHEJ1, NSMCE3, PALB2, RAD51, RAD51C, RFWD3, SLX4, UBE2T, XRCC2)	□ REFLEX to Whole Exome Sequencing**(See additional details below)

**Whole exome sequencing (WES) orders require completion of the WES Test Requisition. Also, inclusion of biological parental samples is strongly encouraged to assist with the analysis of WES and to increase test yield. Please visit our website at www.cincinnatichildrens.org/exome to obtain the required documents. WES testing will NOT be started until all forms are completed and received by the lab.

☐ Spinal Muscular Atrophy - SMN1/SMN2 Copy Number Analysis by MLPA



REQUIRED: Patient Name: Date of Birth:	REQUIRED: Patient Name:	Date of Birth:
--	-------------------------	----------------

TEST(S) REQUESTED CONTINUED

Stickler Syndrome Gene Panel (13 genes)
BMP4, COL11A1, COL11A2, COL2A1, COL9A1, COL9A2, COL9A3, GZF1, LOXL3,
LRP2. PLOD3. SOX9. VCAN

- ☐ Reflex to Cleft and Craniofacial Gene Panel
- ☐ REFLEX to Whole Exome Sequencing** (See additional details below)

\square Treacher Collins Syndrome and Mandibulofacial Dysostos	is Gene Pane
(10 genes) DHODH, EDNRA, EFTUD2, POLR1A, POLR1B, POLR1C,	POLR1D,
SF3B4, TCOF1, TXNL4A	

- ☐ Reflex to Cleft and Craniofacial Gene Panel
- ☐ REFLEX to Whole Exome Sequencing** (See additional details below)

Othe

"Whole exome sequencing (WES) orders require completion of the WES Test Requisition. Also, inclusion of biological parental samples is strongly encouraged to assist with the analysis of WES and to increase test yield. Please visit our website at www.cincinnatichildrens.org/exome to obtain the required documents. WES testing will NOT be started until all forms are completed and received by the lab.

CUSTO	VI CEN	IE CEO	LIENI	CINI	
CUSIU	VI GEN	IE SEG	UEN	CIN	Œ

Gene(s) to be analyzed (specify):

Only genes with clear published functional relationship to rare diseases are accepted.

Suspected syndrome/ condition:

Please choose one of the following:

- ☐ Full gene(s) sequencing
- ☐ Full gene(s) sequencing with reflex to deletion and duplication analysis,

if indicated (please see list of genes available for del/dup at www.cincinnatichildrens.org/deldup)

☐ Familial mutation analysis

Proband's name:

Proband's DOB:

Proband's mutation: __

Patient's relation to proband:

If testing was <u>not</u> performed at CCHMC, please include proband's report and at least 100ng of proband's DNA to use as a positive control.

DELETION AND DUPLICATION ASSAY

Gene(s) to be analyzed (specify):

Please see list of available genes at: www.cincinnatichildrens.org/deldup

Suspected syndrome/ condition:

Please choose one of the following:

- ☐ Deletion and duplication analysis of gene(s) specified above
- ☐ Deletion and duplication analysis of gene(s) specified above with reflex to sequencing, if indicated
- ☐ Analysis of gene(s) specified above from previously analyzed deletion and duplication
- \square Familial deletion analysis

Proband's name: _

Proband's DOB: _

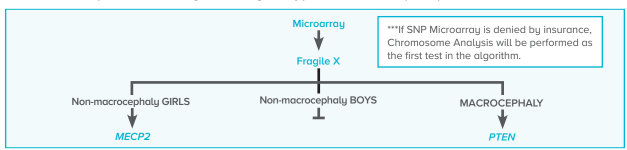
Proband's mutation: _

Patient's relation to proband:

If testing was <u>not</u> performed at CCHMC, please include proband's report and at least 100ng of proband's DNA to use as a positive control.

NEURODEVELOPMENTAL REFLEX GENETIC TESTING ALGORITHM

Tests will be performed sequentially based on the path that your patient follows in the Neurodevelopmental Reflex Test algorithm. Charges will apply to the tests completed in the patient's defined Neurodevelopmental Reflex Test algorithm. Testing will only proceed to the next step if the previous test result is uninformative.



SPECIMEN REQUIREMENTS

- Routine & High Resolution Chromosome Analysis: 3-5 mL blood (NaHep)
- Optical Genome Mapping (OGM) Genome-wide and Targeted Analysis:
 3 mL blood (NaHep) or (EDTA), tissue (1cm x 1cm), or punch biopsy
 (2mm tissue in sterile transport media or saline)
- SNP Microarray:
 - 3 mL blood (NaHep) and 3 mL blood (EDTA)
- FISH Tests: 1-3 mL blood (NaHep)
- Neurodevelopmental Reflex Genetic Testing:
 3 mL blood (NaHep) and 3-5 mL blood (EDTA)
- Fragile X DNA Testing: 3 mL blood (EDTA)
- Fanconi Anemia Chromosome Breakage Study:
 5–10 mL blood (NaHep), 5–10 mL bone marrow (NaHep), or skin biopsy (3-4 mm tissue in sterile transport media)

- ABCD1 del/dup by MLPA, EpiSignature Complete, EpiSignature Targeted, MECP2 del/dup by MLPA, Prader-Willi/Angelman by MLPA, Spinal Muscular Atrophy - SMN1/SMN2 Copy Number Analysis & Deletion/Duplication Assay: 3 mL blood (EDTA)
- ABCD1, FANCA, FANCC, FANCG, MECP2, PTEN & Custom Gene Sequencing:
 3 mL blood (EDTA), saliva collection kit*, or 6 cytobrushes
- Bloom syndrome Sister Chromatid Exchange (SCE) analysis:
 3-5 mL blood (NaHep)
- Cleft and Cranlofacial, Chromosome Breakage Disorders, Fanconi Anemia, Rubinstein-Taybi and Related Syndromes, Stickler Syndrome & Treacher Collins Syndrome and Mandibulofacial Dysostosis Gene Panels:
 - 3 mL blood (EDTA) or saliva collection kit*

*Call the office at 513-636-4474 to obtain saliva collection kits